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Figure 33A is a schematic of the pUA4-4 vector. Also shown in figures 33B-33C is the 5345 bp nucleotide sequence of the vector (SEQ ID NO:17).

On page 6, please replace the paragraph starting on line 7 with the following:

Figure 34A is a schematic of the pTU4 vector. Also shown in figures 34B-34C is the 5337 bp nucleotide sequence of the vector (SEQ ID NO:19).

 \int On page 6, please replace the paragraph starting on line 9 with the following:

Figure 35A is a schematic of the pTT5.14 vector. Also shown in figures 35B-35C is the 5395 bp nucleotide sequence of the vector (SEQ ID NO:21).

On page 6, please replace the paragraph starting on line 11 with the following:

Figure 36A is a schematic of the pTP8-5 vector. Also shown in figures 36B-36C is the 5337 bp nucleotide sequence of the vector (SEQ ID NO:23).

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On page 6, please replace the paragraph starting on line 13 with the following:

Figure 37A is a schematic of the pTP5-1 vector. Also shown in figures 37B-37C is the 5277 bp nucleotide sequence of the vector (SEQ ID NO:25).

On page 6, please replace the paragraph starting on line 15 with the following:

Figure 38A is a schematic of the pTP4a2 vector. Also shown in figures 38B-38C is the 5327 bp nucleotide sequence of the vector (SEQ ID NO:27).

On page 6, please replace the paragraph starting on line 17 with the following:

Figure 39A is a schematic of the pTP3-1 vector. Also shown in figures 39B-39D is the 5338 bp nucleotide sequence of the vector (SEQ ID NO:29).

On page 6, please replace the paragraph starting on line 19 with the following:

Figure 40A is a schematic of the pTU5 vector. Also shown in figures 40B-40H is the 5337 bp nucleotide sequence of the vector (SEQ ID NO:31).

On page 6, please replace the paragraph starting on line 21 with the following:

Figure 41A is a schematic of the pGT6 vector. Also shown in figures 41B-41H is the 4773 bp nucleotide sequence of the vector (SEQ ID NO:32).

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On page 6, please replace the paragraph starting on line 23 with the following:

Figure 42A is a schematic of the pJQ5 vector. Also shown in figures 42B-42I is the 5034 bp nucleotide sequence of the vector (SEQ ID NO:33).

Con page 6, please replace the paragraph starting on line 25 with the following:

Figure 43A is a schematic of the pJO6.1 vector. Also shown in figures 43B-43I is the 4950 bp nucleotide sequence of the vector (SEQ ID NO:34).

 \mathcal{L} On page 6, please replace the paragraph starting on line 27 with the following:

Figure 44A is a schematic of the pJQ4 vector. Also shown in figures 44B-44I is the 4974 bp nucleotide sequence of the vector (SEQ ID NO:35).

On page 6, please replace the paragraph starting on line 29 with the following

Figure 45A is a schematic of the pPQ10.1 vector. Also shown in figures 45B-45H is the 5164 bp nucleotide sequence of the vector (SEQ ID NO:36).

On page 6, please replace the paragraph starting on line 31 with the following:

Figure 46A is a schematic of the pJQ3 vector. Also shown in figures 46B-46I is the 4965 bp nucleotide sequence of the vector (SEQ ID NO:37.

On page 7, please replace the paragraph starting on line 1 with the following:

Figure 47A is a schematic of the pUG4 vector. Also shown in figures 47B-47C is the 5295 bp nucleotide sequence of the vector (SEQ ID NO:38).

On page 7, please replace the paragraph starting on line 3 with the following:

Figure 48A is a schematic of the pUB8.11 vector. Also shown in figures 48B-48I is the 5001 bp nucleotide sequence of the vector (SEQ ID NO:40).

On page 7, please replace the paragraph starting on line 5 with the following:

Figure 49A is a schematic of the pTP11-1 vector. Also shown in figures 49B-49C is the 5387 bp nucleotide sequence of the vector (SEQ ID NO:41).

On page 7, please replace the paragraph starting on line 7 with the following:

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Figures 50A-B illustrate[s] the actin promoter and its corresponding nucleotide sequence (SEQ ID NO:43).

On page 7, please replace the paragraph starting on line 9 with the following:

Figure 51 illustrates the Aleurain-NPIR delete structure. The corresponding nucleotide sequences are also shown (SEQ ID NO:45).

On page 7, please replace the paragraph starting on line 11 with the following:

Figure 52 illustrates the SEE1 (senescence enhanced) promoter sequence (SEQ ID NO:46).

On page 7, please replace the paragraph starting on line 13 with the following: Figure 53 illustrates the SEE1 (senescence enhanced) promoter sequence plus the vacuolar aleurain signal/NPIR sequence (SEQ ID NO:47 and 48).

On page 16, please replace the paragraph starting on line 30 with the following:

The present invention provides for methods of changing the cell wall structure of transgenic plants and therefore, making them more digestible. The method comprises introducing a ferulic acid esterase coding sequence into the cells of a plant. Operably linked to the coding sequence is a promoter that can be either constitutive or inducible and signal sequences that serve to target expression of the coding sequence in the desired organelle in the desired cell of the plant. The signal sequences can be either or both N terminal or C terminal sequences.

On page 21, please replace the paragraph starting on line 27 with the following:

In addition to targeting expression to specific organelles, it may be desirable to retain the expressed FAE in the Golgi or endoplasmic reticulum. The well known ER retention signal, KDEL (SEQ ID NO:97), can be added to the 3' end of the coding polynucleotide.

On page 26, please replace the paragraph starting on line 14 with the following:

A genomic clone for FAE1 (see Figures 1-3, SEQ ID NO:1 and 2) was used as the starting point for the preparation of an intronless FAE1 encoding DNA sequence. The sequence for the genomic clone is given in Figures 2 and 3 (SEQ ID NO:1 and 2). Separate fragments for

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both FAE exons were recovered by PCR from a 5.5kb EcoRI fragment of the genomic clone in pLITMUS28, and 'cDNA' created by overlapping PCR. See Figure 4.

On page 26, please replace the paragraph starting on line 20 with the following:

Two 5' primers were used. FAE-S5 which amplifies the entire reading frame (including the Aspergillus signal), and FAE-N5 which amplifies only the mature protein (i.e. has *no* signal). A number of codons are optimized (underlined in primer sequences below). The overlap product may be derived from either FAE-I5 (wild type, SEQ ID NO:49) or FAE-I3 (conserved Ser changed to Ala, SEQ ID NO:50) primers, allowing production of enzymatically inactive protein to check toxicity. As shown in Figure 5, overlapping of PCR products made with FAE-I5 and FAE-I3 creates two possible uninterrupted reading frames (SEQ ID NO:3 and 4). If the complement to FAE-I5 serves as the template when recombined then the encoded protein retains the serine moiety and the esterase is functional (highlighted serine is at active site, SEQ ID NO:95). If the FAE-I3 primer serves as the template the serine is replaced with an alanine and the esterase is inactivated (highlighted alanine in bottom amino acid sequence given in Figure 5, SEQ ID NO:96).

On page 27, please replace the paragraph starting on line 1 with the following:

Where possible, codon usage has been optimized in constructed reading frames (codon choice based on published barley preferences).

FAE-I5 (SEQ ID NO: 49)

FAE-I3 (SEQ ID NO: 50)

CCGGCCACGCCCTCGGCGCCTCCCTGGCGGCACTC 35-mer

FAE-N5 (SEQ ID NO: 51)

CTAAAGCTTACCATGGCGGCCGCCTCCACGCAGGGCATCTCCGA 44-mer

FAE-S5 (SEQ ID NO: 83)

CTAAAGCTTAACATGAAGCAGTTCTCCGCCAA 32-mer

FAE-3 (SEQ ID NO: 52)

TCTAAGCTTGCGGCCGGCCGGCCAGGTGCATGCGCCGCTCGTCATCCC 50-MER

On page 27, please replace the paragraph starting on line 28 with the following:

The nos terminator from **pMA406** (Ainley & Key (1990) PMB 14:949-60) was amplified by PCR using primers TER5 and TER3 to generate a fragment with the following sequence (SEQ ID NO:53):

(Pst1) (Not 1)

(HindIII) (Xbal)

On page 29, please replace the paragraph starting on line 3 with the following:

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Vector sequences were confirmed by sequencing. Two artifacts were found. Firstly, the redundant codon in TER5 was found to be AAA in one clone, which was subsequently used as the source of all KDEL fusions (ie peptide sequence is KPLKDEL (SEQ ID NO:85), rather than EPLKDEL (SEQ ID NO:86) as designed). See Figure 9. Secondly, an additional base is found at the site of the redundant codon in one clone, creating a frameshifted terminal peptide (ETTEG, Figure 10 SEQ ID NO:87) which was used as a control in some constructs.

On page 30, please replace the paragraph starting on line 18 with the following:

PCR primers

TER-5 (SEQ ID NO:54)

AGACTGCAGACCATGGCGGCCGCGKAACCACTGAAGGATGAGCTGTAAAGAAGCAGATC GTTCAAACATTTG 72-MER (The KDEL stop codon is underlined.)

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TER-NOT (SEQ ID NO:55)

AAGACTGCAGACCATGGCGG 20-MER

TER-3 (SEQ ID NO:56)

AGATCTAGAAGCTTATCGATCTAGTAACATAGATGACACC

ALECUT (SEQ ID NO:57)

CTAGGCGCCGCGCGGGGAGGAGGCGACGCGAC

GLYB (SEQ ID NO:58)

GAGGGTGTATTCGGTATCGAGTTGCAGGTTCGTATC



GLY3 (SEQ ID NO:59)

CTCGATACCCATTACACCCTCACGCCTTTCGA

On page 30, please replace the paragraph starting on line 38 with the following:

i. Rice actin promoter and 1st intron

Initial vectors (Figures 11 and 12) were constructed from pCOR105 which was subsequently found to contain a 5bp deletion relative to the published sequence which destroys the AccI site (GTAGGTAGAC, SEQ ID NO:60, deleted bases underlined) and may affect splicing at the adjacent 3' site. The original rice actin sequence in this region (GTAGGTAG, SEQ ID NO:84) was therefore restored using oligonucleotide NCO-ACT (CTCACCATGGTAAGCTTCTACC TACAAAAAAGCTCCGCA, SEQ ID NO:61) by replacing the BgIII/HindIII fragment with a PCR product, to produce vector pPQ10.1.

On page 32, please replace the paragraph starting on line 13 with the following:

PCR Primers

SEE-VAC (SEQ ID NO: 62)

AACCATGGCGGCCGCGCGCTCGGTGACGGGCCGGAT

SEE-NCO (SEQ ID NO: 63)

TTCGGTACCATGGCCAGGTATAATTATGG

SEE-ATG (SEQ ID NO: 64)

CTGCGCCGGCGAGATGGMCGTGCACAAGGAG

On page 32, please replace the paragraph starting on line 30 with the following:

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This is from the original clone and has the peptide sequence:

MKQFSAKHVLAVVVTAGHALAASTQGI (SEQ ID NO:88).

On page 33, please replace the paragraph starting on line 1 with the following:

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Peptide sequence is MAAASTQGI (SEQ ID NO:89) (underlined motif is common to all constructs). Truncation of the signal sequence in (a) above was carried out by PCR with mutagenic primer FAE-N5.

On page 33, please replace the paragraph starting on line 6 with the following:

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The barley aleurain vacuolar signal sequence (See Figure 13; Swissprot database accession number P05167, SEQ ID NO:10) was derived entirely from overlapping primers

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(ALE-5, ALE-3, ALE-CUT ALE-CAP-5 and ALE CAP-3). Following primer annealing at 37°C and extension with T4 DNA polymerase in the presence of dNTPs according to manufacturers instructions, PCR with flanking primers ALE-5 and ALE-3 was carried out. The product was 'polished' with T4 DNA polymerase, purified, digested with NotI and cloned into EcoRV/NotI digested pCOR105-nos terminator vector (see above). ALE-3 contains redundancies so that clones encoding NPIR or NPGR motifs may be recovered. Two versions of the signal, with and without the vacuole targeting motif, were produced, to give putative vacuolar NPIR and apoplast (NPGR) signal sequences.

On page 33, please replace the paragraph starting on line 18 with the following:

PCR Primers

ALE-5 (SEQ ID NO: 65)

GGAATTCGTAGACAAGCTTACMATGGCCCACGCCCGCGTCCT 41-MER

ALE-3 (SEQ ID NO: 66)

TATCCATGGCGCCGCGCGGTCGGTGACGGCCGGMYCGGGTTGGAGTCGGCGAA

55-MER

ALE-CUT (SEQ ID NO: 67)

CTAGGCGCCGCGGGAGGAGGCGACGGCGAC 33-mer

ALECAP-5 (SEQ ID NO: 68)

GCGACGGCGACGCCGTGGCCAGCACGCGCGAGCGCCAGGAGGACGCCG

54-MER

ALECAP-3 (SEQ ID NO: 69)

TCGCCGTCGCCTCCTCCTCCTTCGCCGACT 33-MER

On page 34, please replace the paragraph starting on line 5 with the following:

R36

A Golgi targeting vector, <u>pJQ3.2</u>, was made by inserting a reading frame encoding the relevant rat sialyl transferase (RST) motif (See Figure 14, SEQ ID NO:11_and 12. RST motif shown to function in plants by Boevink P, Oparka K, Cruz SS, Martin B, Betteridge A, Hawes C, (1998) PLANT JOURNAL <u>15</u> 441-447 Stacks on tracks: the plant Golgi apparatus traffics on an actin/ER network) into vector pPQ10.1, and replacing the EcoRI/NotI promoter/signal fragment of **pJO6.3** with the fragment from this vector. Briefly, the RST motif was constructed by annealing oligonucleotides RST-F1A, RST-F1B, RST-F2A and RST-F2B, and amplifying the product with RST-5AD and RST-3A. This product was cloned and sequenced. Clones were

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found to have a deletion which was corrected by PCR with RST-RPT, followed by overlap-PCR and cloning of products.

On page 34, please replace the paragraph starting on line 17 with the following:

PCR primers

RST-5AD (SEQ ID NO: 70)

ACTAAGCTTAAGGAGATATAACAATGATCCACACCAACCTCAA

RST-F1A (SEQ ID NO: 71)

TTCCATGATCCACACCAACCTCAAAAAGAAGTTCTCCCTCTTCAT

RST-F1B (SEQ ID NO: 72)

RST-F2A (**SEQ ID NO: 73**)

A

TATAGATCTGCGTGTGGAAGAAGGGCTCCGACTACGAGGCCCTCCAAGCCAAGG

RST-F2B (SEQ ID NO: 74)

CATTTGGAACTCCTTGGCTTGGAGGGTG

RST-3A (SEQ ID NO: 75)

AACCATGGCGGCCGCCATTTGGAACTCCTTGGCT

RST-RPT (SEQ ID NO: 76)

TATAGATCTGCGTGTGGAAGAAGGGCTCCGACTACGAGGCCCTCCCAAGCCAAGG

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On page 35, please replace the paragraph starting on line 14 with the following:

PCR primers

PPI-AP1 (SEQ ID NO: 77)

GGAATTCGTAGACAAGCTTACMATGGMCGTGCACAAGGAGGT

PPI-AP2 (SEQ ID NO: 78)

GATCAGGAGGTAGGCWACGAAGTTWACCTCCTTGTGC

PPI-AP3 (**SEQ ID NO: 79**)

CCTACCTCCTGATCGTSCTCGGCCTCCTCTTGCTCGT

PPI-AP4 (SEQ ID NO: 80)

CCTTGGCGTCCACGTGCTCCATGGCGGAWACGAGCAAGAGGAG

PPI-AP5 (SEQ ID NO: 81)

GTGGACGCCAAGGCCTGCACCCKCGAGTGCGGCAACCTC







PPI-AP6 (SEQ ID NO: 82)

GGAATTCGCGGCCGGGCAGATGCCGAAGCCGAGGTTGCCGCACT

On page 35, please replace the paragraph starting on line 31 with the following:

This was derived directly from the genomic clone (see Example 1) as a Nco1-Sph1 fragment (Sph end filled with T4 polymerase) which replaces the Nco1-Not1 region of a standard actin -FAE vector (Not1 end filled with T4 DNA polymerase).

Expression vector linker alone [CTW-PVAAA, SEQ ID NO:93] (plant optimised C-terminus for vacuole, golgi and apoplast vectors).

On page 36, please replace the paragraph starting on line 5 with the following:

CTW is the peptide sequence of the Aspergillus FAE COOH end and is here provided by oligo FAE3. In this primer the reading frame is extended to provide the additional amino acids PVAAA (SEQ ID NO:91) which are partially encoded by the Not1 site used for cloning downstream signals see c) and d) below. Some COOH amino acids /motifs may affect compartment targeting, the PVAAA (SEQ ID NO:91) sequences are expected to be neutral in this respect while the native Aspergillus end may not be.

On page 36, please replace the paragraph starting on line 12 with the following:

(c) Linker plus KPLKDEL (SEQ ID NO:90) [first K is primer artifact, intended to be E] {ER retention vectors)

On page 36, please replace the paragraph starting on line 27 with the following:

The linker used in the above C-terminal targeting sequences was PVAAA (SEQ ID NO:91).

On page 37, please replace the paragraph starting on line 22 with the following:

PCR primer

ALE-G (SEQ ID NO:92)

TATCCATGGCGGCCGGGTCGGTGACGGCCCGGGCTTGGAGTCGGCGAA

In the claims:

The plant of claim 3, wherein the polynucleotide further comprises a polynucleotide that encodes CTWPVAAA (SEQ ID NO:93) at the 3' end.